## Introduction to Plant Breeding

#### A Parochial view

Origins of crops
Scientific approaches 1850... present
Plant & animal breeding compared
Achievements & questions



Matthew 7:18-7:20 A good tree cannot bring forth evil fruit, neither can a corrupt tree bring forth good fruit. Every tree that bringeth not forth good fruit is hewn down, and cast into the fire. Wherefore by their fruits ye shall know them.

## The Scientific approach to plant breeding

#### Two strands:

#### 1. Mendelian:

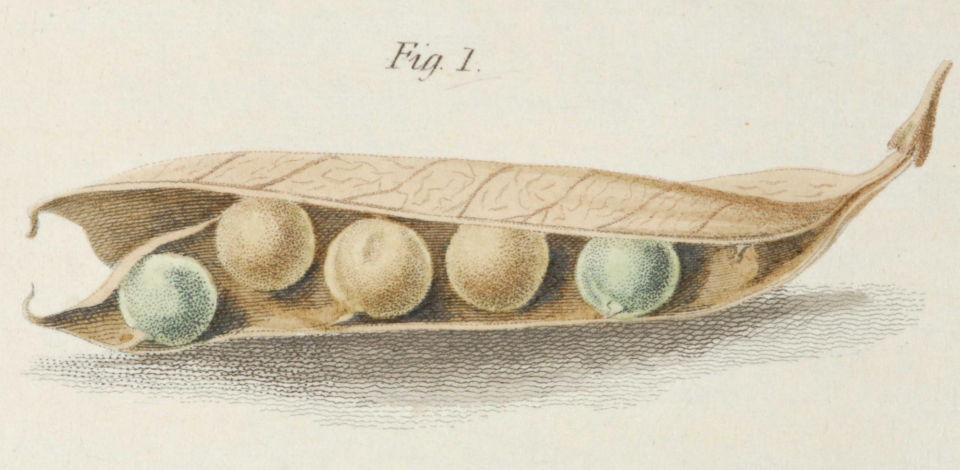
Incorporate information from genes into selection decisions championed by plant breeders

#### 2. Biometric:

Incorporate information from relatives into selection decisions championed by animal breeders

Prospects: we now have the technology to combine the two.

HORT: TRANS. VOLV. PL.9



John Goss (1824) On Variation in the Colour of Peas, occasioned by Cross Impregnation Horticultural Transactions (Series 1) Vol.5, p. 234-237 + 1 fig

#### Some milestones in Mendelian genetics & breeding

1823:	Knight: Dominance,	recessiveness,	and segregation	observed in peas

- 1900: Rediscovery and verification of Mendel's principles
- 1903: Biffen: resistance to stripe rust of wheat is Mendelian recessive.
- 1908: Nilsson-Ehle: seed colour in wheat is due to 3 Mendelian factors.
- 1923: Sax: linkage between quantitative and qualitative traits in beans.
- 1956: Flor: gene for gene hypothesis for host-parasite resistance
- 1965-70 Borlaug: Green Revolution (India & Pakistan) based on dwarfing genes.
- 1983: Beckmann & Soller: RFLPs for genome wide QTL detection and breeding
- 2001: Meuwissen *et al* : Genomic selection proposed

## Wheat Genetic history: plant breeding.



Dwarfing genes reduced

Dwarfing genes reduced

Dwarfing genes reduced

Evaluation of the solution of the solu

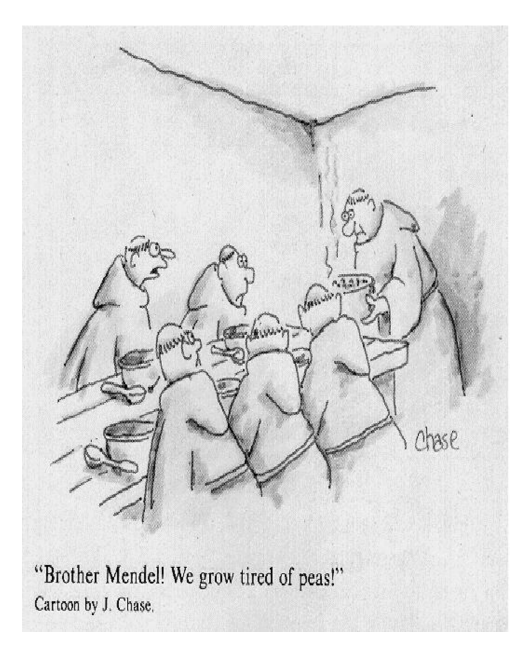
Whigheincreased is susceptibility to biseddition the linesed it with the sewer ing general power and per ear.

• Fungicide



Quantitative methods in plant breeding – March '09

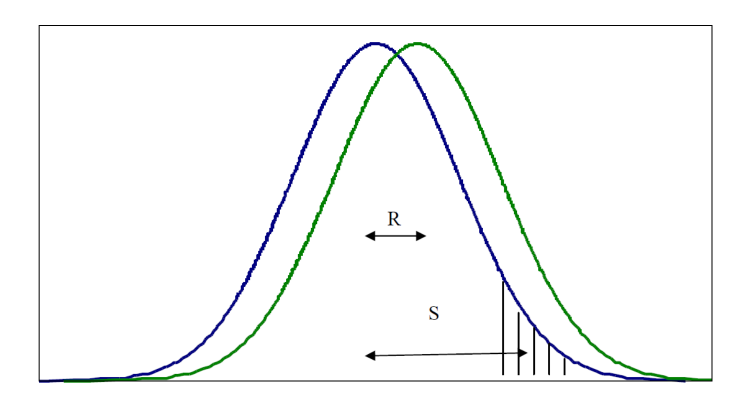
## Information from genes.



## Some milestones in biometrical genetics & breeding

1840-50	de Vilmorin: progeny test in wheat, oat, and sugar-beet breeding.
1889	Galton: publishes Natural Inheritance, a statistical statement of the relative influence of parents
1921	Wright: relationships between relatives
1936	Smith: selection index
1947	Lush: Family merit & individual merit as a basis for selection
1953	Henderson: origins of BLUP
1971	Patterson & Thompson REML
2001	Meuwissen et al: Genomic selection proposed

Both approaches are linked by the breeders' equation  $R = h^2S$ .



Everything in plant (and animal) breeding can be judged by its effect on "the breeders' equation."

The breeders' equation  $R = h^2S$ .

standardized as:

$$R = i h \sigma_g / time / f$$

#### Some arbitrary dates in plants breeding methods

1840-50 de Vilmorin progeny testing

1909 Nilsson-Ehle scientific wheat breeding:

pedigree breeding, bulk breeding

1878-81 Beal corn hybrids yield more

1909 Shull: use of F1 hybrids between inbreds in corn breeding

1924 Blakeslee & Belling report doubled haploids

1939: Golden single seed descent

1936 ? haploids and polyploids

#### Some features of plant breeding methods

Replicate genotypes: clones

inbred lines

**DH** lines

F1 hybrids

Heritabilities vary through replication

Inbreeding is quick self: S1,S2..Sn, doubled haploids

Mating systems: selfing, outcrossing

gms, cms, S alleles, ...

Polyploids: haploids, allopolyploids, autopolyploids

Use of ancestral species eg synthetic wheat

GxE generally larger than in animals

Half sibs have a common female parent

## Methods for selection within crosses

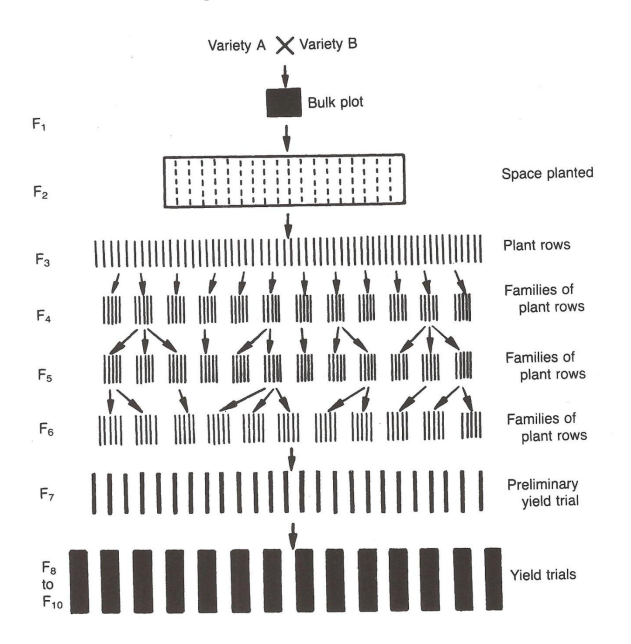
Pedigree breeding

Single seed descent

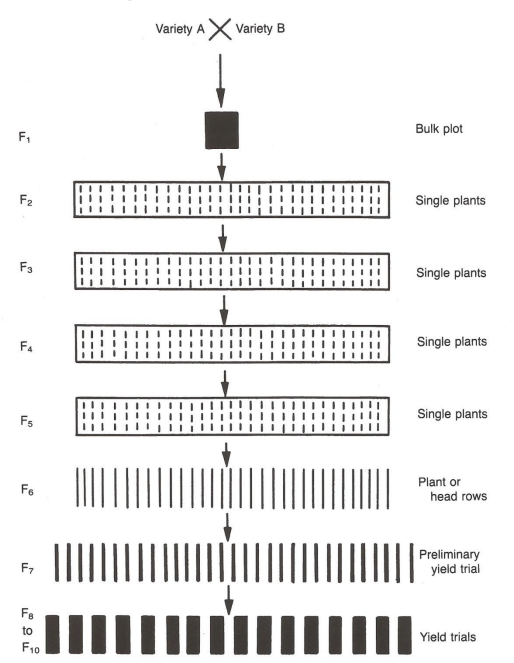
Doubled haploids

Bulk breeding

## Pedigree method



## Single Seed Descent



## Single Seed Descent

Goulden (1939)

Knott & Kumar (1975) wheat

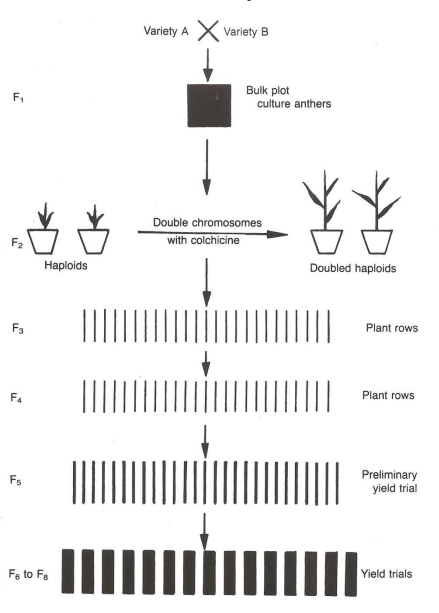
Pedigree breeding: inbreeding & selection concur

SSD: separate inbreeding from selection (faster)

Proposed and developed for breeding.

Use in trait mapping is more recent.

## **Doubled Haploids**



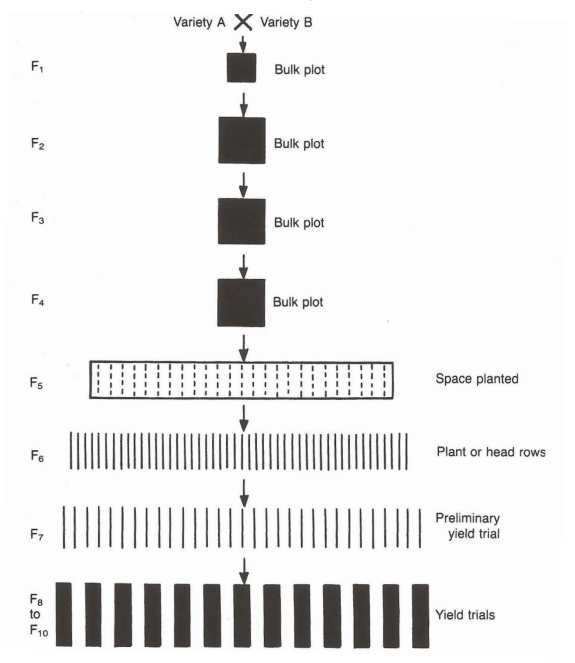
## **Doubled Haploids**

"The practical importance of haploids and polyploids in plant breeding is being quickly recognised and it seems possible that their artificial production will be simply a matter of technique in the near future." *Imperial Bureau of Plant Genetics, 1936* 

Faster than SSD

Expensive
Low efficiency in some crops
Less recombination

## **Bulk Breeding**



## Bulk breeding

As slow as pedigree breeding

Encourage selection in the bulk (natural & artificial)

F2s contribute unequally to inbred lines

Long history (Allard, Harlan)

Not much used in commercial plant breeding.

Regularly rediscovered by academics. And funded!

## Hybrid breeding

General combining ability

Specific combining ability

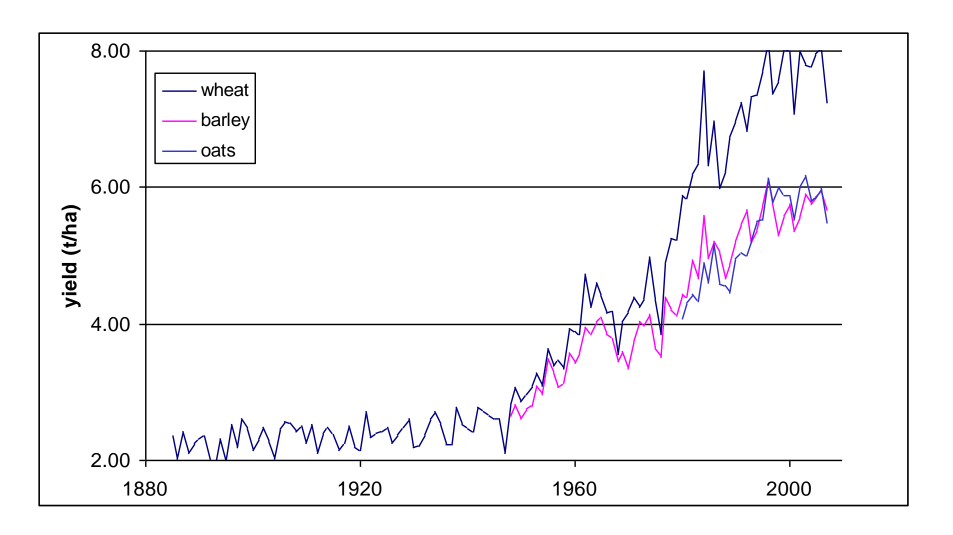
Circulant partial diallels

Heterotic groups

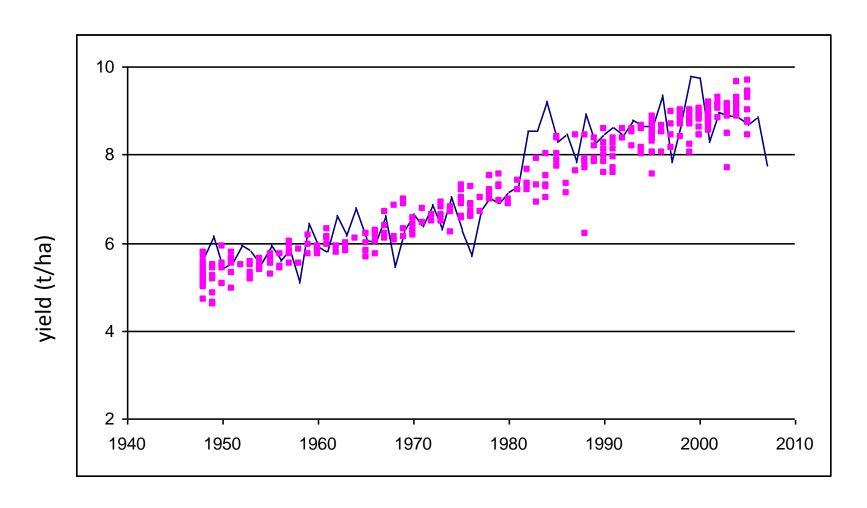
Reciprocal recurrent selection

More money

## Cereal yields in the UK



#### winter wheat genetic and environmental trends

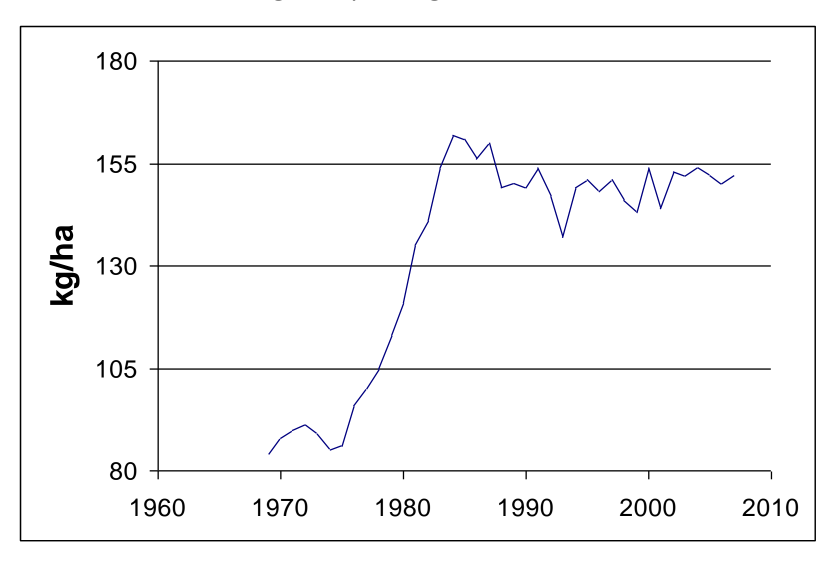


first year in trial

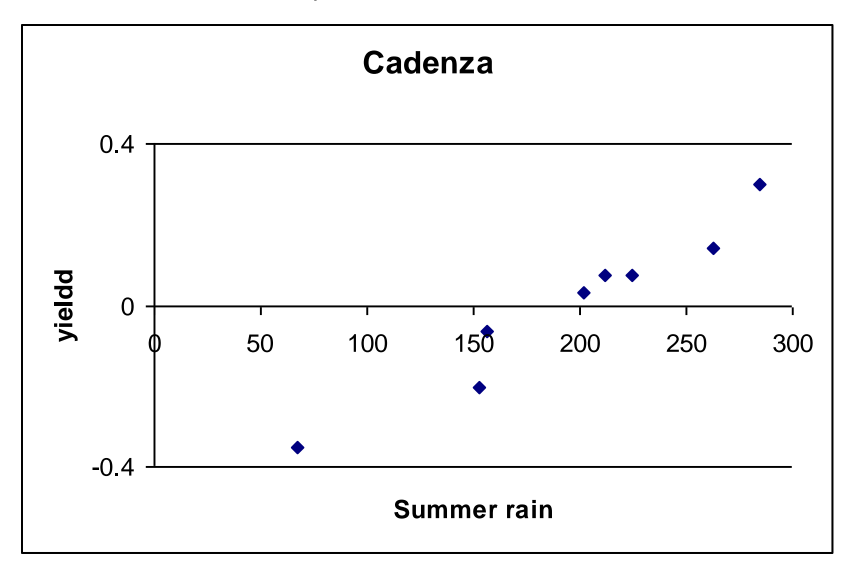
# Linear trends in yield (t/ha) 1982-2007 NL/RL trials

varieties	years
0.074	0.010
0.060	-0.006
0.071	0.010
0.109	0.108
0.105	0.112
0.064	-0.019
	0.074 0.060 0.071 0.109 0.105

## N use for tillage crops: England & Wales



## Screen for sensitivity to climatic stress?



#### Some challenges & questions; a personal view

Have yields stopped rising?

Should we care about GxE?

What proportion of quantitative variation has originated by mutation since domestication: should we sample wild and old germplasm for yield QTL?

Do we get enough recombination?

Why are yield and quality negatively correlated?

Are the days of breeding to exploit natural variation numbered by GM?

What is the best design of a breeding programme to exploit GS?

# WHY HUNT? WHY GATHER? JOIN THE SOLITION!







Build

permanent,

First off — we don't just look around for our food ... we actually grow some of it ourselves, where we live!



Plant and animal domestication is the key. We grow edible plants ourselves, right out of the ground, time after time!

You can start by joining us in the village! Leave your troubles behind!\*





\*Some hunting and gathering may be necessary to maintain dietary variety and avoid famine.

Reshape your environment.

Reshape your environment.

Sociable.

Special offer free booklets. The Pleasures of Porridge Earn Your Animols' Respect

complex,

socielies.

## Your KEYS to a BETTER LIFE!

#### **Harness Plant Power!**

- Learn how the seeds you drop can become next fall's crop!
- Use seed selection to make future plants more productive and easier to harvest!
- Preserve and store surpluses for hard times!
- Invent new ways of preparing and cooking plant foods!

#### Put Animals To Work For You!

- Learn which species are slow and submissive!
- Use food and fences to keep them around!
- · Influence their choice of mates!
- . Breed the best and eat the rest!

Disclaimer: Plant and animal domestication can lead to overpopulation, deforestation, erosion, flooding, desertification, materialism, diminished nutrition, cavities, and television. Caution advised. YOUR RESULTS MAY YARY

CONTRACTOR CONTRACTOR

# Monday pm

 Population genetics and linkage disequilibrium

#### **Population Genetics**

#### **Books**

Felsenstein

http://evolution.genetics.washington.edu/

Weir Genetic Data Analysis 2<sup>nd</sup> ed.

http://statgen.ncsu.edu/powermarker/

## GH Hardy 1877-1947

"There is no permanent place in the world for ugly mathematics."



"I am reluctant to intrude in a discussion concerning matters of which I have no expert knowledge, and I should have expected the very simple point which I wish to make to have been familiar to biologists."

# Hardy-Weinberg Equilibrium 1908

A sufficient condition for no evolution to occur within a Mendelian population is that mutation, selection, and chance effects are all absent and that mating is at random.

The hereditary mechanism, of itself, does not change allele frequencies. The constancy of genotype frequencies then follows from the presence of random mating.

## **Population Genetics**

## The Hardy-Weinberg Law

Nothing changes except for:

mutation selection sampling variation (drift) migration non-random mating

#### **Population Genetics**

#### The Hardy-Weinberg Law

genotype AA Aa aa Χ **2Y** Z frequency all A 1/2 A, 1/2 a all a alleles

Frequency of A gamete  $X + \frac{1}{2} 2Y = p$ say

Frequency of a gamete  $Y + \frac{1}{2}2Y = 1-p = q$  say

with p + q = 1

female gamete (freq) A (p) a (q) male gamete (freq)  $AA(p^2)$ (p) Aa (pq) Α Aa (pq)  $aa (q^2)$ (q) a  $\rightarrow$ 

AA Aa aa  $p^2$  $q^2$ 2pq

 $p^2 + \frac{1}{2} 2pq = p(p+q) = p$ Frequency A:

## **Polyploids**

$$(p_1A_1+p_2A_2+p_3A_3....p_nA_n)^p$$

Eg Bufo pseudoraddei baturae



#### **Population Genetics**

Non-random mating.

AA

Aa

aa

$$p^2 + pqf$$

2pq(1-f)

q<sup>2</sup>+pqf

Selfing series

generation

AA

Aa

aa

0

 $p^2$ 

2pq

 $q^2$ 

1

 $p^2 + pq/2$ 

pq

 $q^2+pq/2$ 

2

 $p^2 + pq3/4pq/2$ 

 $q^2+pq3/4$ 

3

 $p^2 + pq5/8pq/4$ 

= p

 $q^2+pq5/8$ 

 $\infty$ 

p<sup>2</sup>+pq

0

 $q^2+pq=q$ 

#### **Population Genetics**

#### Mixed selfing and random mating

$$f = s/(2-s)$$

where s is the proportion of seed set by selfing or

$$f = (1-t)/(1+t)$$

where t is the proportion of seed set by random mating

#### **Population Genetics**

#### Wahlund effect

Subdivided populations have reduced heterozygosity:

Frequency in population 1 = 
$$p_1 = p+x$$
  
Frequency in population 2 =  $p_2 = p-x$   
Average heterozygosity =  $(2p_1q_1 + 2p_2q_2)/2$   
=  $(p+x)(1-p-x) + (p-x)(1-p+x)$   
=  $2pq - 2x^2$ 

Cross pops– observe excess of hets:

$$(p+x)(1-[p-x]) + (1-p-x)(p-x)$$
  
=  $2pq + 2x^2$ 

Explanation for heterotic pools and composite varieties

#### Linkage Disequilibrium

Random mating between individuals generates equilibrium genotype frequencies at a single locus.

(Hardy-Weinberg equilibrium)

Random assortment of chromosomes in meiosis generates equilibrium frequencies between loci.

(Linkage equilibrium)

At equilibrium:

loc B r(B) s(b)

Loc A

p (A) pr AB ps Ab

q (a) qr aB qs ab

Rearranging:

AB Ab aB ab

pr ps qr qs

Same in the next generation

#### With arbitrary frequencies

Loc A loc B B b
A W X
a y Z

Compare observed and expected with  $\chi^2$ 

AB Ab aB ab Observed W X Z Expected pr ps qr qs O - E+D-D -D +D

D = observed frequency minus expected frequency

AB Ab aB ab pr ps qr qs +D -D +D -D  $D = p(AB) - p(A) \cdot p(B)$ 

or 
$$-D = p(aB) - p(a) \cdot p(B)$$
  
etc.

#### Some properties of the D

Max value is 0.25, when p(A)=p(B)=0.5

At other allele freqs. max. value can be small eg

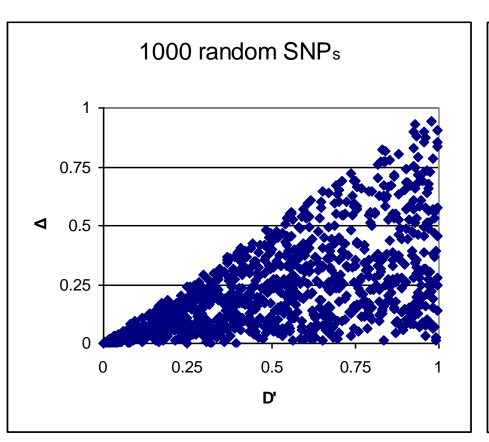
$$p(A)=p(B)=0.9$$
  $D_{max} = 0.09$ 

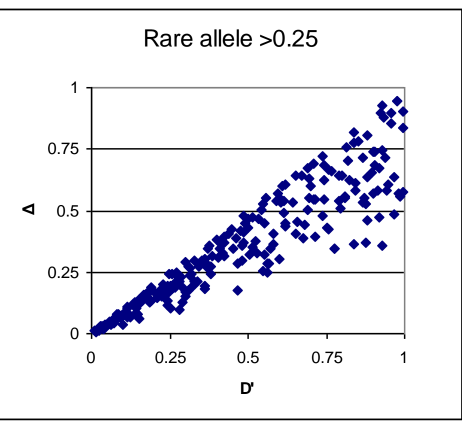
To make interpretation easier, define:

D' = D / D<sub>max</sub> range 0-1  
or
$$\Delta = D range 0-1$$

$$\sqrt{(p(A)p(a)p(B)p(b)}$$

#### Comparison of LD measures





 $\Delta \rightarrow 1$ : allele freqs match, two haplotypes

 $D' \rightarrow 1$ : allele freqs don't matter, three haplotypes

#### LD measures for multiple alleles

Calculate D' or r<sup>2</sup> for each pair of alleles in turn.

Take the average, weighted by the expected frequency  $(p_1p_2)$ 

Estimates tend to be biased upwards in small samples. The bias can be quite large.

Correct by permutation testing.

The decay of Linkage Disequilibrium

$$D_1 = (1-\theta) D_0$$

$$D_t = (1-\theta)^t D_0$$

# gens	unlinked	5cM	0.5cM	I 50k
0	1	1	1	1
1	0.50	0.95	1	1
10	0	0.60	0.95	1
100	0	0.01	0.61	0.95
1000	0	0	0.01	0.61
10000	0	0	0	0.01

# **Proof**

To decay, LD needs recombination. Recombination need double heterozygotes

Arbitrarily select gamete type AB to follow over 1 generation:

$$P_{(AB)} = 2(pr + D)(qs + D) (1-\theta) /2 + 2(ps-D)(qr-D) \theta /2$$
(non recs from AB/ab) (recs from Ab/aB)

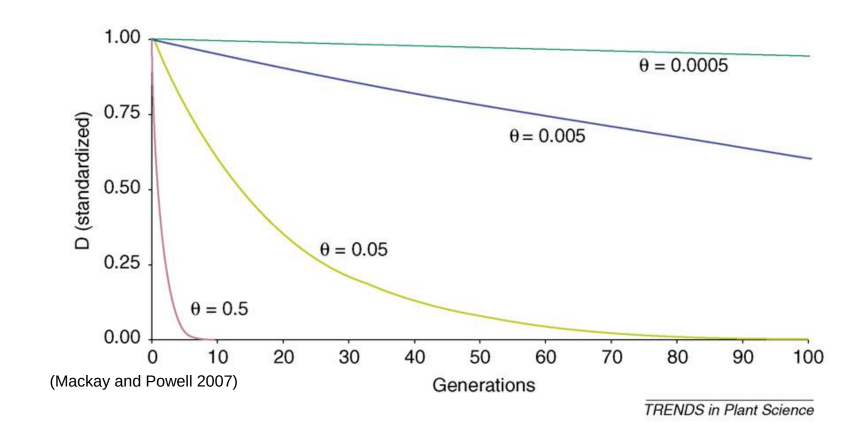
Ignore terms not involving  $\theta$  to get change in  $P_{(AB)}$ 

$$= [-(pr + D)(qs + D) + (ps-D)(qr-D)]\theta = -\theta D$$

New value of D is therefore D –  $\theta$ D = D(1- $\theta$ )

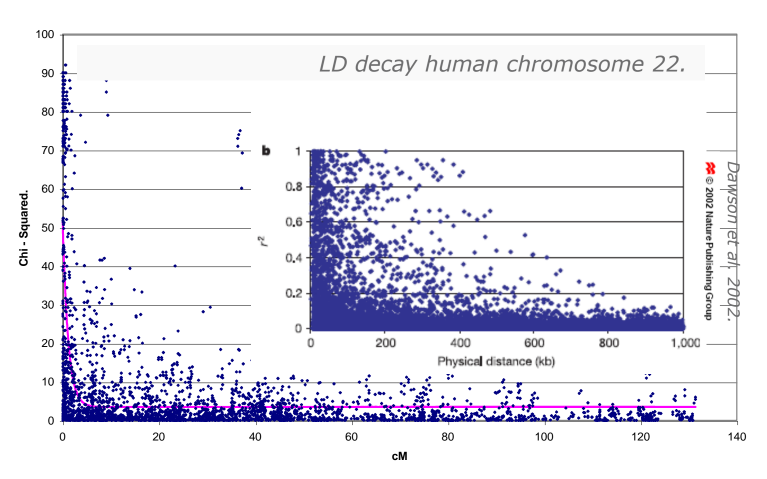
Over t generations:  $D_t = D_0(1 - \theta)^t$ 

### LD decays with time and recombination fraction



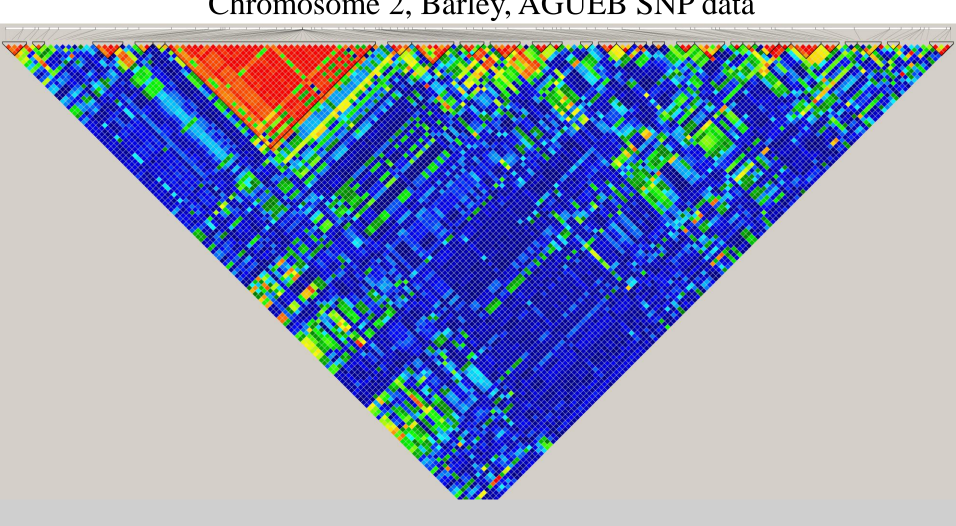
### Decline in LD with genetic distance

Decline of between marker association over genetic distance. UK wheat all genomes.



# LD in Barley varieties

Chromosome 2, Barley, AGUEB SNP data



### The Causes of Linkage Disequilibrium

Mutation

Sampling

drift, founder effect

Migration

Selection

#### Mutation

Gen.	Allele freq	D'	Δ
0	1/2N	1	0
X	?	?	?

Although mutation generates LD, this is not very interesting. It is the fate following mutation which is important.

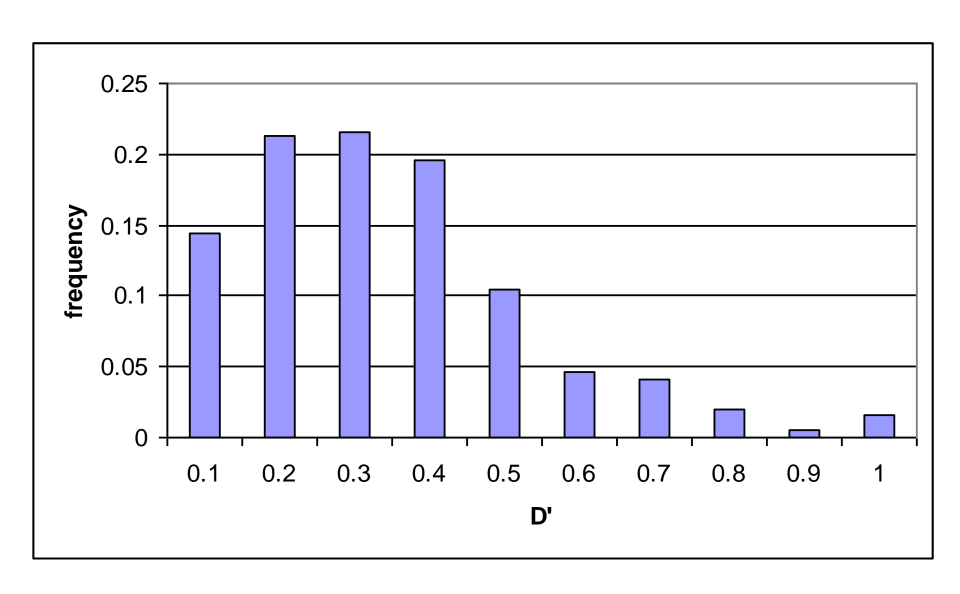
Drift

$$\mathbf{\mathcal{E}}(\Delta^2) = 1$$

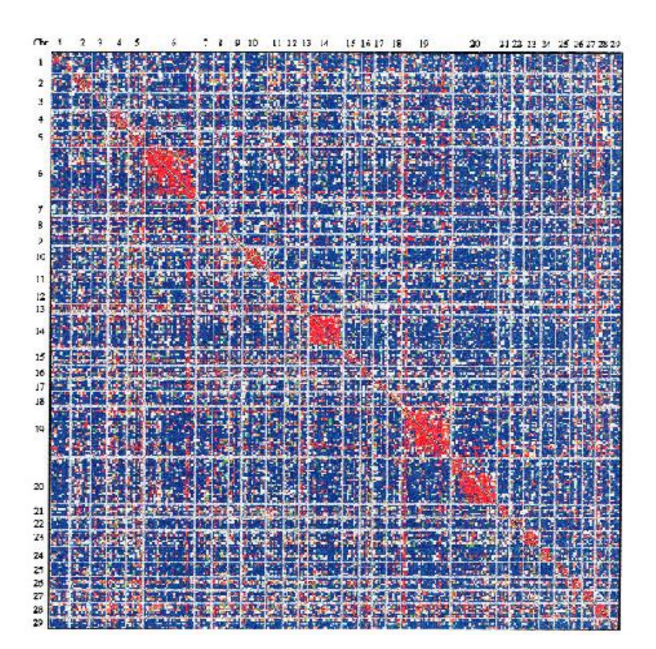
$$1 + 4N_e\theta$$

On average, as population size and recombination increase, LD falls

### Distribution of LD in founder population size 10







### **Migration**

Pop 1 (no LD)

Pop 2 (no LD)

 $p_1r_1$  (AB)

 $p_2r_2$  (AB)

1:1 mix

What is the freq. of AB

Observe

 $\frac{1}{2}(p_1r_1+p_2r_2)$ 

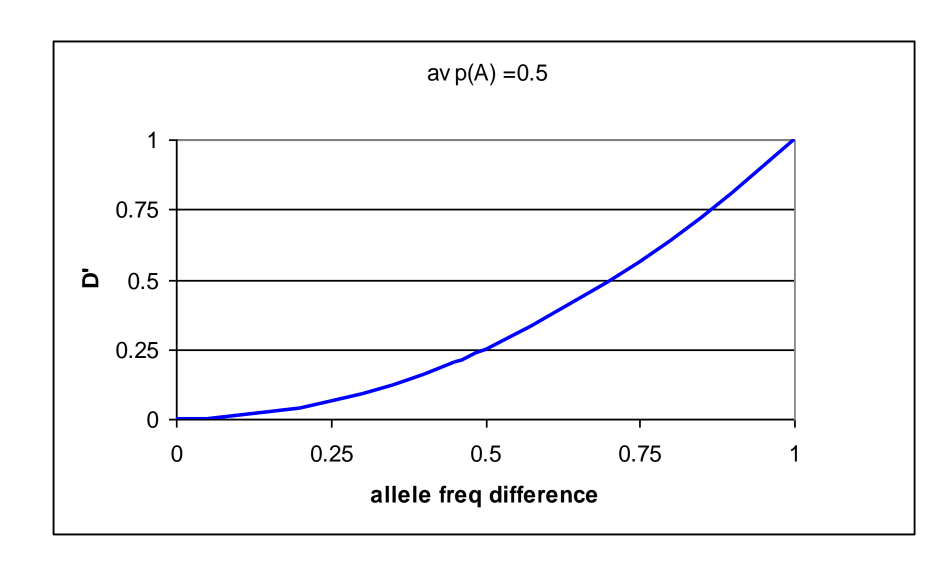
Expect

 $^{1}/_{4} (p_{1}+p_{2})(r_{1}+r_{2})$ 

$$D = \frac{1}{4} (p_1 - p_2)(r_1 - r_2)$$

Zero if 
$$p_1 = p_2$$
 or  $r_1 = r_2$ 

### **Migration – population admixture**



### Hitch-hiking

Allele frequencies change at a locus as a result of selection.

As a result, closely linked polymorphisms change in frequency too.

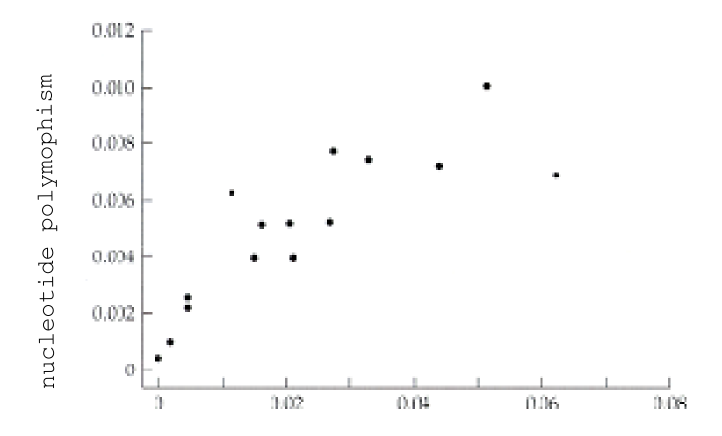
Hitch-hiking generates LD over the whole linked region.

Is important in regions of low recombination.

These are the gene-rich regions – more opportunities for selection.

# Hitch-hiking: evidence from Drosophila





Rate of recombination

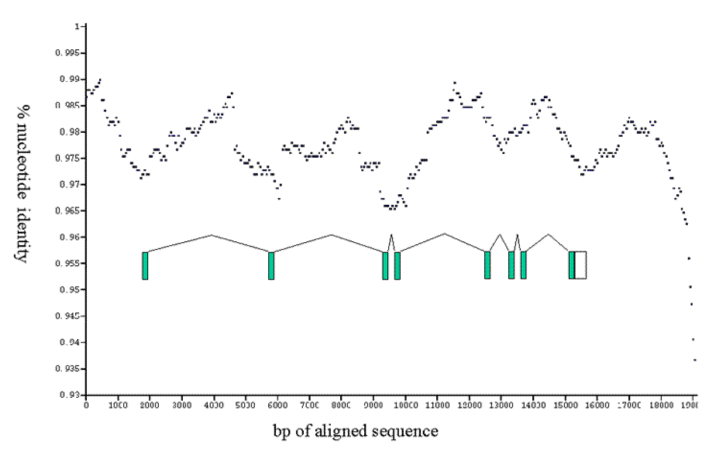
### An example of hitch-hiking in man.

The Morpheus gene family – function unknown – found in a class of segmental duplications.

20x normal rate of amino acid substitution.

Non synonymous substitution rate > synonymous.

Sequence alignment of two human copies of morpheus gene family.



0 16 K bases

### So what?

Deleterious SNPs at a high frequency are likely to be of interest.

One way they may rise in frequency is through hitch-hiking.

Therefore – look for footprints of hitch-hiking:

High LD / low recombination / gene rich regions Lower heterozygosity and freq. of neutral SNPs Higher heterozygosity and freq. of nsSNPs

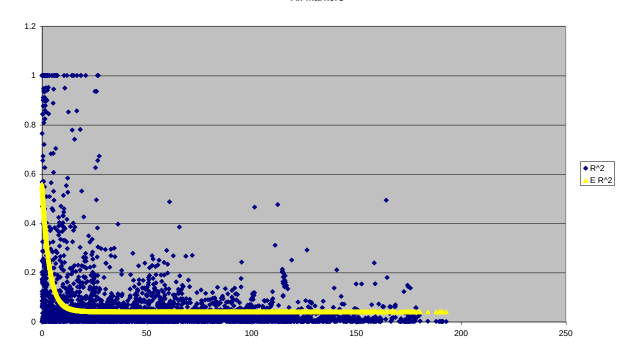
### Plotting and Modelling LD

$$E(\Delta^2) = 1/(1+4N_e\theta)$$

$$E(D') = L + (H-L)(1-\theta)^{t}$$

#### B Genome UK

All Markers



#### Haplotypes

Methods of determining phase:

is AaBB:

<u>AB,</u> <u>ab</u>

or

<u>Ab</u>, <u>aB</u>

Pedigree CEPH families

Sequencing short range

Clarke Algorithm easy to understand

EM much software - snphap

Evolutionary methods Phase